UNCLASSIFIED

AD 274 278

Reproduced by the

ARMED SERVICES TECHNICAL INFORMATION AGENCY
ARLINGTON HALL STATION
ARLINGTON 12, VIRGINIA



UNCLASSIFIED

NOTICE: When government or other drawings, specifications or other data are used for any purpose other than in connection with a definitely related government procurement operation, the U.S. Government thereby incurs no responsibility, nor any obligation whatsoever; and the fact that the Government may have formulated, furnished, or in any way supplied the said drawings, specifications, or other data is not to be regarded by implication or otherwise as in any manner licensing the holder or any other person or corporation, or conveying any rights or permission to manufacture, use or sell any patented invention that may in any way be related thereto.

X-RAY INVESTIGATIONS OF Mn,P, Mn,P and Ni,P

Stig Rundqvist

Institute of Chemistry, University of Uppsala Uppsala, Sweden.

Technical Note No. 30 Contract No. AF 61(052)-40 December 27, 1961

ASTIA

APR 18 1962

TISIA

B

The research reported in this document has been sponsored in part by the

AIR FORCE OFFICE OF SCIENTIFIC RESEARCH

of the AIR RESEARCH AND DEVELOPMENT COMMAND, UNITED STATES AIR

FORCE, through its European Office.

ray investigations of Mn_P, Mn_P, and Ni_P.

Stig Rundqvist

Institute of Chemistry, University of Uppsala, Uppsala, Sweden.

The crystal structures of Mn_3P (Fe₃P type), Mn_2P and Ni_2P (revised C 22 type) have been refined by single-crystal methods. Both Mn_2P and Ni_2P have appreciable homogeneity ranges. X-ray powder investigations in the Mn_2P system indicate that the existence of the phase described as Mn_3P_2 is doubtful.

The crystal structures of transition metal phosphides with the compositions Me₃P and Me₂P have been the subject of earlier investigations by the author 1,2,3,4,5 in order to provide material for detailed crystal-chemical comparisons within this group of compounds. In the present paper, accurate structure determinations of Mn₃P, Mn₂P, and Ni₂P are reported. In addition, some phase-analytical data for the Mn-P system are presented and discussed.

Experimental

Preparation. The starting materials for the preparations were electrolytic manganese (AB Ferrolegeringar, Stockholm, claimed purity higher than 99.9%), nickel rods, spectrographically standardized (Johnson, Matthey & Co, Ltd., London), and red phosphorus (purity higher than 99%). Master alloys were prepared by dropping pellets of red phosphorus into molten metal, contained in crucibles of pure alumina (Degussit Al 23 from

Degussa, Frankfurt, Germany). The melting was done by induction heating under a parified argon atmosphere. Alloys of various compositions were made by mixing portions of the crushed master alloys and heating in evacuated and sealed silica tubes. The nickel phosphides did not attack silica appreciably, even after very long heat-treatments. The manganese phosphides, however, attack silica, and the annealing time for manganese phosphides was therefore kept as short as possible.

X-ray work. The phase analyses were performed by X-ray powder methods only. Powder photographs were recorded in Guinier-type focussing cameras with $Cr\underline{K} \ll 1$ radiation using silicon $(\underline{a} = 5.4305 \ \text{Å})$ as the internal calibration standard. The accuracy of a single lattice parameter measurement is estimated to be 0.04 %, but lattice parameter differences larger than 0.02 % measured for the same phase in different alloys are probably significant.

single-crystal fragments with roughly uniform cross-sections not exceeding 0.05 mm were selected from the crushed alloys.

Weissenberg photographs were taken using zirconium-filtered

Mok radiation. The multiple-film technique, with thin iron

foils between successive films, was used. The intensities were

estimated visually by comparison with calibrated spots. Corrections for Lorentz and polarisation factors, Fourier series

summations, structure factor calculations, and calculations of

interatomic distances were made on the electronic digital computer

BESK with programmes available at BESK. Atomic scattering factors were taken tables given for manganese by Watson and Freeman⁶,

for nickel by Thomas and Umeda ⁷, and for phosphorus by Tomiie

and Stam ⁸. The real part of the dispersion correction for the scattering factors of the atoms as given by Dauben and Templeton ⁹ was inserted for each of the metal atoms in the structure factor calculations. Standard deviations for the atomic positions were estimated using Cruickshank's ¹⁰ equation. (List of observed and calculated structure factors can be obtained from this Institute on request).

The manganese - phosphorus system.

Earlier work on the Mn-P system is summarized by Hansen 11 . X-ray investigations by Arstad and Nowotny 12 showed the existence of Mn₂P, Mn₂P and MnP. Thermal analytical, microscopic, and X-ray investigations by Berak and Heumann 13 confirmed the findings of Arstad and Nowotny, and in addition indicated the existence of Mn₂P₂.

In the present investigation, which is restricted to the range 0-50 atom % phosphorus, the only intermediate phases observed were Mn₃P, Mn₂P and MnP. Within the limits of experimental error, the lattice parameters of Mn₃P and MnP were unchanged in alloys with different compositions and heat-treatments. No appreciable lattice parameter variations for Mn₂P were observed in two-phase Mn₃P + Mn₂P alloys (see Table 1), but variations were observed in alloys containing more than 33 ¹/3 atom % phosphorus. In order to study this effect more closely, a special investigation of the Mn₂P - MnP part of the system was undertaken. Since the results differ from those obtained by Berak and Heumann, the work will be described in some detail.

A large number of alloys with the composition $Mn_{1.5}^P$ were annealed in silica tubes at various temperatures between 800°C

and 1090°C for periods varying from one hour to seven days, and subsequently cooled. The rate of cooling was found to have a marked influence on the final state of the samples, and three different methods of cooling the alloys from the annealing temperature to room temperature were therefore used: 1. cooling in air (room temperature attained in 5-10 minutes), 2. dropping the alloys in water without breaking the silica capsules (room temperature in 10-20 seconds), 3. dropping the alloys in water and breaking the capsules (room temperature in less than 3 seconds). The last method will be referred to as "rapid quenching" in the following text.

The X-ray powder photographs of all these alloys showed that the two phases MnpP and MnP alone were present. The MnP diffraction lines were invariably sharp, whereas the MnoP lines were more or less diffuse, depending on the cooling rate. The diffraction lines of MnoP were quite sharp for rapidly quenched alloys, which had been annealed at temperatures below approximately 1060°C. The unit cell dimensions decreased progressively with increasing annealing temperature (see Table 1). The aircooled alloys gave rather diffuse $\mathrm{Mn}_{\mathcal{O}}\mathrm{P}$ lines. The unit cell dimensions were only slightly smaller than those obtained for Mn_2P in two-phase $Mn_3P + Mn_2P$ alloys. The diffraction lines were generally very broad for alloys which had been quenched in water without breaking the silica capsules. In a few cases, two different sets of $\mathrm{Mn}_2\mathrm{P}$ lines, one sharp and one diffuse, were discernible in the powder photographs of these alloys. For each sharp line, a corresponding diffuse line with a slightly smaller diffraction angle was observed. In some instances, a similar effect was also noticed for rapidly quenched alloys.

which had been annealed at temperatures above 1060°C.

All these observations consistently indicate that MnoP has a finite homogenity range, the phosphorus-rich limit of which moves with increasing temperature towards the phosphorusrich side of the diagram. MnP is precipitated during the cooling of Mn. 5P alloys, while the MnoP phase becomes more metal-rich. In rapidly quenched alloys, the secondary precipitation of MnP is negligible. The powder photographs of these alloys exhibit sharp Mn_oP lines. This indicates a well-crystallized condition of the phase, which has the phosphorus-rich limiting composition corresponding to the annealing temperature used. For lower rates of cooling, the secondary precipitation of MnP is appreciable. The broadening of the MnoP diffraction lines is attributable to coring effects. In extreme cases, the supercooling may produce the double set of MnoP lines referred to above. The set of sharp lines belongs to phosphorus-rich material retained un-decomposed on quenching, while the set of diffuse lines corresponds to more metal-rich MnoP which is formed at lower temperatures. The powder photographs exhibit all types of line intermediate between the extremes of very broad lines on the one hand and two separate sets on the other. There is therefore no reason to believe that the double set of lines indicates a two-phase region, in which equilibrium exists between two Mn₂P-type phases of different composition.

As mentioned above, the only phases observed in the range $33^{-1}/3 - 50$ atom % P are Mn_2P and MnP. This is in contrast to the report by Berak and Heumann $^{-13}$ (B & H), who claim the existence of an intermediate phase with the composition Mn_3P_2 . This phase is stated to form peritectically at $1090^{\circ}C$ (only $5^{\circ}C$ above

the eutectic temperature) and decompose eutectoidally at 1002°C. The failure to detect any Mn₃P₂ phase in the present investigation may be due to unsuccessful annealing and/or quenching techniques. It is felt, however, that the arguments presented by B & H for the existence of Mn₃P₂ are not convincing. An attempt to reconcile their observations with the present findings can be made as follows.

The thermal arrest at 1002°C observed by B & H (the somewhat scattered experimental points were obtained only on cooling) may be explained by thermal effects, possibly obscured by supercooling, which would accompany the secondary precipitation of MnP from the phosphorus-rich MnoP phase. The microscopic examination by B & H of an $Mn_{1.5}^{P}$ alloy, quenched from 1080°C, showed a homogeneous phase together with traces of MnoP and MnP. The homogeneous phase, claimed to be the new Mn_3P_2 phase, may in fact have been phosphorus-rich MnoP, which had been retained undercomposed on quenching. The powder photograph of a further $Mn_{1.5}P$ alloy, quenched from $1080^{\circ}C$, was stated to contain new lines in addition to the MnoP pattern. Since the authors do not claim any high accuracy for their X-ray measurements, and no X-ray data for the new phase were given, it is not unreasonable to believe that their powder photograph contained a double set of MnoP lines, possibly in addition to MnP lines, mentioned before, powder photographs of this type have been obtained by the present author.

The homogeneity range of NioP.

In a previous communication ¹⁴ on the nickel-phosphorus system the phase $\mathrm{Ni}_{12}\mathrm{P}_5$ was described. In the present investigation it was found that there exists a two-phase region $\mathrm{Ni}_{12}\mathrm{P}_5 + \mathrm{Ni}_2\mathrm{P}$ up to at least $1000^{\circ}\mathrm{C}$. The lattice parameters of $\mathrm{Ni}_2\mathrm{P}$ were measured in two-phase $\mathrm{Ni}_{12}\mathrm{P}_5 + \mathrm{Ni}_2\mathrm{P}$ alloys quenched from various temperatures up to $1000^{\circ}\mathrm{C}$. The measured unit cell dimensions dit not show detectable variations and were $\underline{a} = 5.865$ Å, $\underline{c} = 3.387$ Å. These values are significantly larger than those obtained for the single-phase $\mathrm{Ni}_2\mathrm{P}$ sample used for the single-crystal work (see below). The unit cell dimensions for $\mathrm{Ni}_2\mathrm{P}$ in the last-mentioned sample were $\underline{a} = 5.859$ Å, $\underline{c} = 3.382$ Å. $\mathrm{Ni}_2\mathrm{P}$ accordingly exhibits contraction of the unit cell with increasing phosphorus content analoguous to that found for $\mathrm{Mn}_2\mathrm{P}$. The equilibria between $\mathrm{Ni}_2\mathrm{P}$ and more phosphorus-rich intermediate phases are not yet known, and were not studied in the present investigation.

The structure determination of Mn_P.

According to Årstad and Nowotny 12 , Mn₃P is isostructural with Fe₃P. This is confirmed by the present investigation. Weissenberg photographs of Mn₃P were taken with crystals rotated about the <u>a</u> and <u>c</u> axes. The structure was refined from successive electron density maps and difference maps projected on the <u>a c</u> and <u>a b</u> planes. For the 129 F (<u>h k</u> 0) values observed, the final R-value of 0.064 was obtained. An overall, isotropic temperature factor with $\underline{B} = 0.44_9$ \underline{A}^2 was applied. The corresponding figures for the 92 F (<u>h</u> 0 1)-values observed were $\underline{R} = 0.048$, $\underline{B} = 0.34_5$ \underline{A}^2 .

(The very strong (004) reflection was omitted on account of extinction). For each atom, two sets of \underline{x} and \underline{y} parameters were obtained, one from the $\underline{\beta}(\underline{x},\underline{y})$ and one from the $\underline{\beta}(\underline{x},\underline{y})$ projection. The final differences between these parameter values did not exceed 0.0005 for any atom. The parameter values quoted below are weighted averages, obtained by giving the values from the centrosymmetric $\underline{x},\underline{y}$ projection twice the weight of those from the noncentrosymmetric $\underline{x},\underline{y}$ projection.

Final structure data for Mn3Pare as follows:

Space group
$$\underline{I} \stackrel{-}{4} - (\underline{S}_{4}), \underline{Z} = 8$$

 $\underline{a} = 9.181 \text{ Å; } \underline{c} = 4.568 \text{ Å; } \underline{U} = 385.0 \text{ Å.}$

Atoms in	8(<u>g</u>)	<u>x</u>	$Q(\overline{\mathbf{x}})$	¥	δ(<u>y</u>)	<u>z</u>	♂(<u>z</u>)
Mn _I		0.0807	0.0002	0.1071	0.0002	0.2279	0.0006
$^{\mathtt{Mn}}$ II		0 .3 567	0.0002	0.0319	0.0002	0.9863	0.0006
Mn		01721	0.0002	0.2192	0.0002	0.7531	0.0006
P		0.2935	0.0004	0.0450	0.0004	0.4880	0.0011

Interatomic distances are listed in Table 2.

The structure determinations of Mn_2P and Ni_2P .

Single-crystals of Mn_2P were selected from a two-phase $Mn_3P + Mn_2P$ alloy. Good single-crystals of Ni_2P proved to be more difficult to obtain, since they were found to be very sensitive to mechanical deformation, which resulted in the X-ray patterns

exhibiting elongated and diffuse spots. It was found, however, that crystal fragments picked from crushed alloys gave single-crystals suitable for X-ray work after a 30 minute heat-treatment at 930°C.

The hexagonal $\operatorname{Mn_2P}$ and $\operatorname{Ni_2P}$ structures both belong to the revised \underline{C} 22 ($\operatorname{Fe_2P}$) structure type 1 . The space group is \underline{P} 6 2 \underline{m} with three metal atoms in 3 (\underline{f}), three metal atoms in 3 (\underline{g}), two phosphorus atoms in 2 (\underline{e}) and one phosphorus atom in 1 (\underline{b}). Both structures were refined by successive electron density projections on the basal plane. During the refinements it was found that the temperature factor of the 3 (\underline{g}) metal atoms was much higher than that of the 3 (\underline{f}) atoms. Individual isotropic temperature factors were therefore introduced, which lowered the \underline{R} -value for $\operatorname{Mn_2P}$ from 0.058 to 0.045 for the 67 \underline{F} (\underline{h} \underline{k} 0)-values observed, and for $\operatorname{Ni_2P}$ from 0.075 to 0.063 for the 51 \underline{F} (\underline{h} \underline{k} 0)-values observed. Electron counts did not indicate different scattering parameters for the two sets of metal atoms.

Final structure data for Mn₂P are as follows:

Space group
$$\underline{P} \stackrel{-}{6} 2 \underline{m} - (\underline{D} \frac{3}{3\underline{h}}); \underline{Z} = 3$$

 $\underline{a} = 6.081 \text{ Å}; \underline{c} = 3.460 \text{ Å}; \underline{U} = 110.8 \text{ Å}^3$

	x	$\delta(\underline{x})$	$\underline{\mathtt{B}}$ (\mathtt{A}^2)
$3 \text{ Mn}_{I} \text{in } 3 \left(\underline{\mathbf{f}}\right)$	0.2546	0.0004	0.23 ₄
3 Mn _{II} in 3 (<u>g</u>)	0.5943	0.0004	0.430
$2 P_{I}$ in $2 (\underline{e})$			0.40
1 P _{II} in 1 (<u>b</u>)			0.40

Final data for NipP are as follows:

$$\underline{a} = 5.859 \text{ Å; } \underline{c} = 3.382 \text{ Å; } \underline{U} = 100.5 \text{ Å}^3$$

		<u>x</u>	$\mathcal{S}(\underline{\mathbf{x}})$	<u>B</u> (Å ²)
3 Ni _I	in 3 $(\underline{\mathbf{f}})$	0.2575	0.0004	0.32
3 Ni	in 3 (g)	0.5957	0.0005	0,69
2 P _I	i n 2 (<u>c</u>)			0.42
1 P _{II}	in 1 (<u>b</u>)			0.42

The unit cell dimensions for $\mathrm{Mn_2P}$ and $\mathrm{Ni_2P}$ quoted above were obtained from powder photographs of the alloys from which the single-crystal specimen were selected.

Interatomic distances are given in Table. 3. A description of the \underline{C} 22 structure is given in ref. 1.

Concluding remarks

The substantial homogeneity ranges of Mn₂P and Ni₂P have escaped the notice of previous investigators. In the present author's opinion, the results of Berak and Heumann in the Mn₂P - MnP part of the Mn-P equilibrium diagram may be explained simply by the extended Mn₂P single-phase field without the introduction of a new intermediate phase. Clearly, an accurate redetermination of the thermal data for the Mn-P system is desirable.

Extended homogeneity ranges seem to be common in Me₂P-type phosphides. According to Haughton ¹⁵, secondary precipitation of

FeP in two-phase $\text{Fe}_{2}P$ + FeP alloys indicates a wider homogeneity range for Fe,P at higher temperatures. The existence of an extended homogeneity range for Co₂P has been demonstrated by Nowotny 16 and by the present author 2, and there are also signs of a similar effect for RupP. Variations of the FepP unit cell dimensions have not been measured, but for Mn2P, Ni2P, and Co2P the unit cell volumes decrease with increasing atomic ratios of phosphorus/ /metal. For Co₂P it has been shown 2 that the homogeneity range extends from the stoichiometric composition towards the phosphorusrich side of the diagram. This is probably due to a varying number of vacancies on one of the crystallographically non-equivalent cobalt atom sites (Co_{TT}). The structure of Co₂P is closely related to that of Mn₂P, Fe₂P, and Ni₂P, and the coordination of the ${
m Co}_{
m II}$ atoms is closely similar to that of the ${
m Mn}_{
m II}$, ${
m Fe}_{
m II}$ and ${
m Ni}_{
m II}$ atoms. (For more detailed descriptions of these structures, see refs. 1, 2). It is noteworthy that the temperature factors for the $\mathrm{Mn}_{\mathrm{TT}}$ and $\mathrm{Ni}_{\mathrm{TT}}$ atoms are substantially larger than those for Mn_{T} and $\mathrm{Ni}_{\mathrm{T}}.$ The origin of this effect is not easy to explain. However, the effect may indicate that the $\mathrm{Mn}_{\mathrm{II}}$ and $\mathrm{Ni}_{\mathrm{II}}$ atomic positions are more likely to be the sites of vacancies or disorder in non-stoichiometric Mn₂P and Ni₂P.

Acknowledgements

This work has been financially supported by the Swedish State Council of Technical Research and by the Air Force Office of Scientific Research of the Air Research and Development Command, United States Air Force through its European Office under Contract No. AF 61 (052)-40. Facilities for use of the electronic digital computer BESK were given by the Swedish Board for Computing Machinery.

The author wishes to thank Professor G. Hägg for his kind interest. Thanks are also due to Mr. O. Olovsson for excellent assistance with numerical computations.

References:

1.	Rundqvist, S. and Jellinek, F.	Acta Chem. Scand	• <u>13</u> (<u>1959</u>) 425.
2.	Rundqvist, S.	Ibid.	<u>14</u> (<u>1960</u>)1961.
3.	Rundqvist, S.	Ibid.	<u>14</u> (<u>1960</u>)2246.
4.	Rundqvist, S.	Ibid.	<u>15</u> (<u>1961</u>) 342.
5.	Rundqvist, S.	Ibid.	(in print)
6.	Watson, R.E. and Freeman, A.I.	Acta Cryst.	<u>14</u> (<u>1961</u>) 27.
7.	Thomas, L.H. and Umeda, K.	J.Chem.Phys.	<u>26</u> (<u>1957</u>) 293.
8.	Tomile, Y. and Stam, C.H.	Acta Cryst.	<u>11</u> (<u>1958</u>) 126.
9.	Dauben, C.H. and Templeton, D.	H. Ibid.	<u>8</u> (<u>1955</u>) 841.
10.	Cruickshank, D.W.J.	Ibid.	<u>2 (1949)</u> 65.
11.	Hansen, M. Constitution of	Binary All cys, Mc	Graw-Hill,
	New York-Toronto	London, 1958.	
12.	Årstad, O. and Nowotny, H. Z.	physik.Chem.	<u>B38</u> (<u>1938</u>) 356.
13.	Berak, J. and Heumann, Th. Z.	<u>Metallkunde</u>	<u>41</u> (<u>1950</u>) 19.
14.	Rundqvist, S. and Larsson, E.	Acta Chem.Scand.	<u>13</u> (<u>1959</u>) 551.
15.	Haughton, J.L. J. Iro	n Steel Inst.	<u>115</u> (<u>1927</u>) 417.
16.	Nowotny, H. Z. and	rg. Chem.	<u>254 (1947</u>) 31.

Table 1. Lattice parameters (in Å) of Mn₂P in two-phase

Mn₂P + Mn₂P and Mn₂P + MnP alloys, quenched from various temperatures.

Temp.	Two-phase Mn	P+Mn ₂ P alloy	Two-phase Mn ₂ P+MnP alloy		
°C <u>a</u>		<u>c</u>	<u>a</u>	<u>c</u>	
900	6.081	3.460	6.074	3.451	
1000	6.081	3.460	6.068	3.446	
1070	6.080	3. 458	6.059	3.440	

Table. 2. Interatomic distances in Mn₃P (Å)

(Distances shorter than 3.7 Å listed)

	Mn _I	Mn _{II}	Mn _{III}	P
MnI	2.46 ₂ ,2.71 ₄ (2),	2.69 ₈ ,2.84 ₉ ,	2.54 ₃ ,2.74 ₂ ,2.77 ₆ ,	2•35 ₇ •2•43 ₈ •
	3.03 ₅ (2)	3.56 ₅	2.81 ₉ ,2.86 ₃	3•56 ₉
Mn _I i	2.69 ₈ ,2.84 ₉ ,	2.69 ₆ ,2.88 ₀ (2)	2.55 ₃ ,2.60 ₃ ,	2•35 ₂ ,2•36 ₁ ,
	3.56 ₅	3.07 ₂ (2)	2.63 ₉ ,3.54 ₉	2•36 ₇ ,2•37 ₀
Mn _{III}	2.5 ⁴ 3,2.7 ⁴ 2, 2.77 ₆ ,2.81 ₉ , 2.86 ₃	2.55 ₃ ,2.60 ₃ , 2.63 ₉ ,3.54 ₉	2•75 ₄ (2)	2.29 ₅ ,2.37 ₇ , 2.43 ₇ ,3.62 ₄
P	2.35 ₇ ,2.43 ₈ ,	2.35 ₂ ,2.36 ₁ ,	2.29 ₅ ,2.37 ₇	3.50 ₁ (2)
	3.56 ₉	2.36 ₇ ,2.37 ₀	2.43 ₇ ,3.62 ₄	3.64 ₁ (2)

Table 3. Interatomic distances in Mn₂P and Ni₂P (Å)

(Distances shorter than 3.7 Å listed)

Type of distance	Number of equivalent distances	Mn ₂ P	Ni ₂ P	Type of distance	Number of equivalent distances	Mn ⁵ b	N1 ₂ P
Me _I -Me _I	2	2.68 ₂	2.61 ₃ 3.38 ₂	P _I -Me _I	3	2 .3 0 ₄	2.209
Me _I -Me _{II}	2	2.694	2,605	P _I -Me _{II}	6	2•53 ₀	2.456
Me _I -P _I	2	2.76 ₇	2.678	P _I -P _I	2	3.46 ₀	3.38 ₂
Me _I -P _{II}	2	2.322	2.266		3	3•51 ₁	3.383
Me _{II} -Me _{II}	2	3.19 ₉ 3.46 ₀	3.08 ₆ 3.38 ₂	P _{II} -Me _I	6	2.322	2.266
Me _{II} -P _I	4	2.53 ₀	2.456	P _{II} -Me _{II}	3	2.46 ₇ 3.61 ₄	2.36 ₉ 3.49 ₀
Me _{II} -P _{II}	1	2.46 ₇ 3.61 ₄	2.36 ₉ 3.49 ₀	P _{II} -P _{II}	2		3.382